



**PTO/SB/08a**  
**Substitute for Form 1449A/PTO**

## **INFORMATION DISCLOSURE STATEMENT BY APPLICANT**

(Use as many sheets as necessary)

PTO/SB/08a Substitute for Form 1449A/PTO		Application Number	10/565,366
<b>INFORMATION DISCLOSURE STATEMENT BY APPLICANT</b> <i>(Use as many sheets as necessary)</i>		Filing Date	January 23, 2006
		First Named Inventor	HARTUNG, Rolf
		Art Unit	1764
		Examiner Name	To be assigned
		Sheet	1
		Attorney Docket	7601/84486

## U.S. PATENT DOCUMENTS

## FOREIGN PATENT DOCUMENTS

Examiner Initials*	Cite No. <sup>1</sup>	Foreign Patent Document	Publication Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages or Relevant Figures Appear	T
		Country Code <sup>3</sup> -Number <sup>4</sup> -Kind Code <sup>5</sup>				
	B1	EP 0 427 939	05-22-1991	American Cyanamid Co.		
	B2	EP 0 438 311	07-24-1991	Merck & Co. Inc.		
	B3	EP 0 823 416	08-07-1996	Ajinomoto Co., Inc.		
	B4	WO 91/07430	05-30-1991	The Upjohn Co.		

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<sup>1</sup> Applicant's unique citation designation number (optional). <sup>2</sup> See Kind Codes of USPTO Patent Documents at [www.uspto.gov](http://www.uspto.gov) or MPEP 901.04. <sup>3</sup> Enter Office that issued the document, by the two-letter code (WIPO Standard ST.3). <sup>4</sup> For Japanese patent documents, the indication of the year of the reign of the Emperor must precede the serial number of the patent document. <sup>5</sup> Kind of document by the appropriate symbols as indicated on the document under WIPO Standard ST.16 if possible. <sup>6</sup> Applicant is to place a check mark here if English language Translation is attached.

PTO/SB/08b Substitute for Form 1449B/PTO				Application Number	10/565,366
<b>INFORMATION DISCLOSURE STATEMENT BY APPLICANT</b> <i>(Use as many sheets as necessary)</i>				Filing Date	January 23, 2006
				First Named Inventor	HARTUNG, Rolf
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				Examiner Name	To be assigned
				Sheet	2

NON PATENT LITERATURE DOCUMENTS					
Examiner Initials	Cite No. <sup>1</sup>	Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published.			T <sup>2</sup>
	C1	International Search Report for PCT/EP2004/006654 filed June 19, 2004.			
	C2	International Preliminary Report on Patentability for PCT/EP2004/006654 filed June 19, 2004.			
	C3	ALEXANDER, <i>et al.</i> , "A Diastereoselective Synthesis of (2S, 3R, 4S)-2-Amino-1-cyclohexyl-6-methylheptane-3,4-diol, The Abbott Aminodiol," <i>Tetrahedron Letters</i> 37:1961-1964 (1996).			
	C4	BIRCH, <i>et al.</i> , "The Structure and Some Reactions of the Iron Tricarbonyl Complex of Thebaine," <i>J. Chem. Soc. C.</i> 531 (1968).			
	C5	BLÁHA, <i>et al.</i> , "Stereoisomeric Chiral 2,9-Diazabicyclo[4.4.0.]Decane-3-,10-Diones as Models of Dipeptide Grouping: Synthesis, X-Ray, IR, NMR, and CD Studies," <i>Coll. Czech. Chem. Commun.</i> 49:712-742 (1984).			
	C6	CLINGMAN, <i>et al.</i> , "Effect of Amines on Hydrogenolysis of Alkylphenols," <i>J. Org. Chem.</i> 23:276-280 (February 1958).			
	C7	CORRINGER, <i>et al.</i> , "CCK-B Agonist of Antagonist Activities of Structurally Hindered and Peptidase-Resistant Boc-CCK <sub>4</sub> Derivatives," <i>J. Med. Chem.</i> 36:166-172 (1993).			
	C8	DEVANT, <i>et al.</i> , "Steroselektive Aldolreaktion Mit Chiralen Sekundären Acetamiden," <i>Chem. Ber.</i> 119:2191-2207 (1986).			
	C9	EISLER, <i>et al.</i> , "Amino Acids and Peptides. LXV. Analogues of Oxytocin," <i>Coll. Czech. Chem. Commun.</i> 31:4563-4580 (1966).			
	C10	FAUSTINI, <i>et al.</i> , "Stereospecificity in the Transformation of $\alpha$ -Aminoacids into Fluroracids," <i>Tetrahedron Letters</i> 22:4533-4536 (1981).			
	C11	HAYASHI, <i>et al.</i> , "Chiral ( $\beta$ -Aminoalkyl)Phosphines. Highly Efficient Phosphine Ligands for Catalytic Asymmetric Grignard Cross-Coupling," <i>J. Org. Chem.</i> 48:2195-2202 (1983).			
	C12	HARRIS, <i>et al.</i> , "Structure of Ristocetin A: Configurational Studies of the Peptide," <i>J. Am. Chem. Soc.</i> 104: 363-365 (1982).			

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Sheet 3	of 3	Attorney Docket 7601/84486		

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	C13	HOEKSTRA, <i>et al.</i> , "Large-Scale Synthesis of Anticoagulant Decapeptide MDL 28050," <i>Tetrahedron</i> 48:307-318 (1992).		
	C14	ISHIDA, <i>et al.</i> , "Micropeptins 88-A to 88-F, Chymotrypsin Inhibitors from the Cyanobacterium <i>Microcystis aeruginosa</i> (NIES-88)," <i>Tetrahedron</i> 54: 5545-5556 (1998).		
	C15	MINNAARD, <i>et al.</i> , "Synthesis of Enantiomerically Pure Cyclohexylglycine," <i>Synthetic Communications</i> 29(24): 4327-4332 (1999).		
	C16	PLATA, <i>et al.</i> , "The Stereospecific Preparation of an Hydroxyethylene Isotere Precursor via a Novel Piperidine-2,5-Dione Template," <i>Tetrahedron Letters</i> 32(30): 3623-3626 (1991).		
	C17	SCHUDA, <i>et al.</i> , "A Short and Efficient Synthesis of (3S, 4S)-4-[( <i>tert</i> -Butyloxycarbonyl)amino]-5-cyclohexyl-3-hydroxypentanoic Acid Ethyl Ester," <i>J. Org. Chem.</i> 53: 873-875 (1988).		
	C18	TAMURA, <i>et al.</i> , "Guanylpiridine Peptidomimetics: Potent and Selective bis-Cation Inhibitors of Factor Xa," <i>Bioorg. Med. Chem. Lett.</i> 10(8): 745-749 (April 2000).		
	C19	TAMURA, <i>et al.</i> , "A Synthesis of Optically Active $\alpha$ -Cyclohexylglycine," <i>Synthetic Communications</i> 8(5): 345-351 (1978).		

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